



The program FEMB users manual

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Risø - M - 1929	Title and author(s) The Program FEMB Users Manual by Ib Misfeldt	Date March 1977
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		Group's own registration number(s)
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	Abstract This report contains a short description of the two-dimensional diffusion theory finite element program FEMB. Input and output explanations are given together with two examples on the use.	Copies to
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SHORT DESCRIPTION

FEMB is a finite element program for approximate solution of the diffusion-theory-equation

$$(1) \quad + \nabla(D_g(\underline{r}) \nabla \phi_g(\underline{r})) + \Sigma_{r,g}(\underline{r}) \phi_g(\underline{r}) + \\ + \sum_{g' \neq g} \Sigma_{s,g \leftarrow g'}(\underline{r}) \phi_{g'}(\underline{r}) + \frac{1}{k_{eff}} \cdot \chi_g(\underline{r}) \sum_{g'} (\nu \Sigma_f)_{g'}(\underline{r}) \phi_{g'}(\underline{r}) = 0$$

within a domain Ω in the xy-plane with the boundary conditions

$$(2) \quad - D_g \frac{\partial \phi_g(\underline{r})}{\partial n} = \sum_{g'} \gamma_{gg'}(\underline{r}) \phi_{g'}(\underline{r}) - \phi_g(\underline{r})$$

or

$$(3) \quad \frac{\frac{\partial \phi_g(\underline{r})}{\partial n}}{\phi_g(\underline{r})} = L_g(\underline{r})$$

Ω is divided in rectangular elements. Within each element the flux ($\phi_g(\underline{r})$) is approximated by a polynomial of order m.

For more details about the method and the approximation used see ref. 2-4.

The equations are solved for one group at a time using the ordinary power iteration technique. The equation for one group is solved by a pointwise successive overrelaxation technique. There is only one inner iteration for each outer.

The iterations are accelerated by extrapolation for removal of the dominating error-mode.

The iterations are stopped when the local error is less than a given input parameter (eps).

An approximate upper limit for the local error ($\epsilon_{1,max}$) is calculated in this way:

$$\epsilon_{1,max} \leq \frac{\max_i |\phi_i^k - \phi_i^{k-1}|}{\max_i \phi_i^k} \cdot \text{extpl}$$

where k gives the outer iteration number, i runs over all flux points in all groups and extpl is the maximal of the last three extra polation factors used for extrapolation.

APPLICATION

On the whole the program adheres to the input/output conventions of the Rise program TWODIM. It is therefore easy to use for users accustomed to this program.

The program gives very reliable results within reasonable calculation times, but it is not a fast program and should therefore mostly be used where high precision is needed.

The typical precision of a calculation with the program can be seen in ref. 1 and ref. 2.

INPUT SPECIFICATION

As mentioned before there is only little difference between FEMB and TWODIM in the input, therefore the same name for input variables with the same meaning is used.

First comes a complete list of the input parameters in the order in which they appear.

FF:
 Problem nb: } administration input
 day, month, year: }

CMX: number of coarse mesh in the x-direction.

CMY: number of coarse mesh in the y-direction.

NCP: number of compositions=number of ordinary materials + number of boundary conditions.

NB: number of boundary conditions.

NDB: number of diagonal boundary conditions; condition (3)

NG: number of groups.

NRR: number of reaction-rates, for each reaction-rate the program calculates for each element

$$\sum_{g'} (RR_{g', \text{element}} \cdot \int_{\text{element}} \phi_{g'}(\underline{r}) d\underline{r})$$

KDS: if KDS=1 then the reaction-integrals from the last calculation are printed in condensed form as specified by meshx and meshy.

if KDS=2 then the reaction-integrals from all calculations are printed in the same condensed form, as specified by meshx and meshy.

if KDS=0 meshx and meshy are not read.

- NREC: the calculations are repeated NREC times, between each calculation the mesh is divided as specified by FACX and FACY.
- M: gives the order of the interpolation polynomial, for practical reasons M should be less than or equal to 6.
- MX: the maximal number of fine mesh in the x-direction.
- MY: the maximal number of fine mesh in the y-direction.
- XC: array (0: CMX), containing the coordinates of the coarse mesh division in the x-direction.
- YC: as XC.
- FMX: array (1: CMX), gives the number of fine mesh into which each coarse mesh is divided before the first calculation.
- FMY: as FMX.
- FACX: array (1: CMX) containing the number of fine mesh into which each mesh is divided before the next recalculation.
- FACY: as FACX.
- CMP: array (0: CMX + 1, 0: CMY + 1) containing lay-out of composition numbers. The numbers are read as if they form part of the first quadrant in the xy-plane. Each logical record corresponds to one mesh in the y-direction, the first and the last are boundary conditions.

Diffusion data for the NCP compositions:

K: composition number.

For $K \leq NB$ and $K \leq NDB$:

EXTPL: array (1:NG) current/flux ratios for the NG groups at the boundary characterized by the composition number K.

For $NDB < K \leq NB$

GAMMA: array (1:NG, 1:NG) corresponding to boundary condition (2); non-negative diagonal elements, non-positive off diagonal elements and

$$\sum_g \gamma_{gg} \geq 0 \text{ for all } g.$$

For $NB < K \leq NCP$:

Data for the ordinary material, characterized by the composition number K.

D: array (1:NG) diffusion coefficient for the groups.

SS: the modified scattering matrix

$$SS(g, g) = \Sigma_{r, g} = \Sigma_{abs, g} + \sum_{g' \neq g} \Sigma_{s, g' \leftarrow g}$$

$$SS(g, g') = \Sigma_{s, g \leftarrow g'}$$

FS: array (1:NG) normalised fission spectrum.

NFS: array (1:NG), $\sqrt{\Sigma_{fiss}}$.

RR: array (1:NRR, 1:NG) reaction rate cross section, each reaction rate is one logical record, if NRR=0 RR is not read.

MESHX: array (0:MX) the first and the last number in the array must always be 1. All the numbers correspond to a fine mesh interface, if the number is 0 the two adjacent mesh are condensed in the output from the reaction integral.

if KDS=2 MESHX is $(0; \sum_{i=1}^{CMX} FMX(i))$ and the numbers corresponds to the mesh after the first subdivision as specified by FMX

MESHY: as MESHX.

The last cards contain four control variables for each calculation, if -1 is given for the first calculation then default values are used, if -1 is given in the following cards then unchanged values are used for the corresponding variable.

MAXANTALITERATIONER: the maximal number of iterations, default is 50.

EPS: The iterations are stopped when the local error ($\xi_{1, max}$, calculated as described on page 1) is less than eps, default is 0.0001.

EPSMY: Extrapolation is performed when the relative change in the extrapolation factor for two consecutive iterations is less than EPSMY, default is 0.1.

OMEGA: overrelaxation factor, default is 1.2.

The input is now concluded with the administration input cards FF and -1 or a new problem.

Quick reference input specification

Each line in the following corresponds to one logical record and must start on a new card.

Repeated equally structured data is shown as [data] · N where N gives the number of repeats.

```

FF
problem
day, month, year
cmx, cmy, ncp, nb, ndb, ng, nrr, kds, nrec, m
mx, my
xc(0), xc(1), ....., xc(cmx)
fmx(1), fmx(2), ....., fmx(cmx)
facx(1), facx(2), ....., facx(cmx)
yc(0), yc(1), ....., yc(cmy)
fmy(1), fmy(2), ....., fmy(cmy)
facy(1), facy(2), ....., facy(cmy)
cmp(0, cmy+1), cmp(1, cmy+1), ....., cmp(cmx+1, cmy+1)
cmp(0, cmy), cmp(1, cmy), ....., cmp(cmx+1, cmy)

```

$$\text{cmp}(0,0), \text{cmp}(1,0), \dots, \text{cmp}(\text{cmx}+1, 0)$$
$$\begin{bmatrix} k \\ \text{extpl}(1), \text{extpl}(2), \dots, \text{extpl}(ng) \end{bmatrix} \cdot \text{ndb}; \text{lsk} \leq \text{ndb}$$
$$\begin{bmatrix} k \\ \text{gamma}(1,1), \text{gamma}(1,2), \dots, \text{gamma}(ng,ng) \end{bmatrix} \cdot (no-ndb); \quad ndb \leq k \leq nb$$

k		$nb \leq k \leq ncp$
$d(1), d(2), \dots, d(ng)$		
$ss(1,1), ss(1,2), \dots, ss(1,ng), \dots, ss(ng, ng)$		
$fs(1), fs(2), \dots, fs(ng)$		
$nfs(1), nfs(2), \dots, nfs(ng)$		
$rr(1,1), rr(1,2), \dots, rr(1,ng)$		$\cdot (ncp - nb)$
$rr(2,1), rr(2,2), \dots, rr(2,ng)$		
\cdot		
\cdot		
\cdot		
$rr(nrr,1), rr(nrr,2), \dots, rr(nrr, ng)$		

$$\left. \begin{array}{l} \text{meshx}(0), \text{meshx}(1), \dots, \text{meshx}(\text{mx}) \\ \text{meshy}(0), \text{meshy}(1), \dots, \text{meshy}(\text{my}) \end{array} \right\} \text{ if kds} = 1$$

$$\left. \begin{array}{l} \text{meshx}(0), \text{meshx}(1), \dots, \text{meshx}(\sum_{i=1}^{\text{CMX}} \text{FMX}(i)) \\ \text{meshy}(0), \text{meshy}(1), \dots, \text{meshy}(\sum_{i=1}^{\text{CMX}} \text{FMY}(i)) \end{array} \right\} \text{ if kds} = 2$$

if kds = 0 no reading of meshx and meshy

[maxantaliterationer, eps, epsmy, omega] · (nrec+1); -1 ⇒ default
 FF
 -1 (or a new problem) } administration input

OUTPUT DESCRIPTION

Each logical page starts with the usual ADM. heading and can consist of several physical pages.

Page 1 is a listing of input in the order in which it is read, one logical record is read at a time and immediately written.

Page 2 starts with a listing of the parameters used for the following iteration (maxantaliterationer, eps, epsmy, omega). Then comes a listing of some control variables of minor interest for the ordinary user, they are:

OANTAL: number of compositions = NCP.

RTYPER: number of boundaries = NB.

GANTAL: number of groups.

EANTAL: number of elements = number of meshes with ordinary materials.

RANTAL: number of external boundaries.

ANTALUBK: number of unknowns per group.

MAXHUD: the amount of space used for the equations.

ANTALBLK: the number of blocks used for the equations on secondary storage. maxhud/900 gives the minimum number of blocks, if antalblk is much greater, contact the person responsible for the program.

The maximum allowed number of blocks is 600/ng, if more space is needed contact responsible engineer.

After these control variables comes the progress report for the iteration.

IT.NR: the number of this iteration

RESIDUE: $\frac{\max_i |\phi_i^k - \phi_i^{k-1}|}{\max_i \phi_i^k}$; k gives iteration number, i runs over all flux-points in all groups

RATIO: $\frac{\sum_i |\phi_i^{k-1} - \phi_i^{k-2}| / \max_i \phi_i^k}{\sum_i |\phi_i^{k-1} - \phi_i^{k-2}|} \cdot \text{sign}(\phi_j^k - \phi_j^{k-1}) \cdot \text{sign}(\phi_j^{k-1} - \phi_j^{k-2})$;

$$|\phi_j^{k-1} - \phi_j^{k-2}| = \max_i |\phi_i^{k-1} - \phi_i^{k-2}|$$

LAMBDA: $1/k_{\text{eff}}$

KEFF: k_{eff}

DELTAKEFF: $(k_{\text{eff}}^k - k_{\text{eff}}^{k-1})$

TOTALKILDE: $\sum_e \sum_g \left(\int_e \phi_g (\nabla \Sigma_f)_g d\Omega \right)$

e runs over all elements

g runs over all groups

Every time an extrapolation is performed MY and eksfac is written

MY: =RATIO

EKSFAC: $1/(1-MY)$

Page 3 The result of the calculation, k_{eff} and fluxdistribution normalised so that $\phi_{\text{max}} = 9999$

Page 4 average flux for each group with the normalisation mentioned above, and a description of where the maximum occurs.

Page 5 if NRR $\neq 0$ page 5 contains a map for each reaction-rate, the normalisation is done so that the average is 1000 for each reaction integral.

Page 6 if NREC > 0 the following 4 pages contain the same as page 2-5.

EXAMPLE

As a simple example we take the reactor shown in fig. 1.

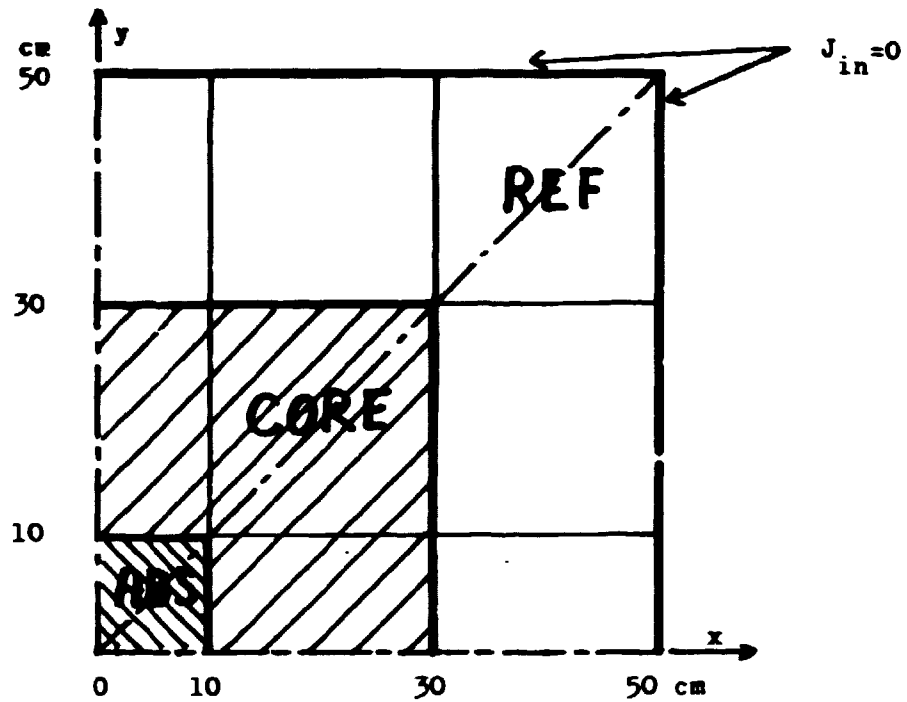


Fig. 1. Geometry specification.

For this reactor we perform a two-group calculation with the following coefficients.

Region	D_1	D_2	$\Sigma_{1 \rightarrow 2}$	Σ_{a1}	Σ_{a2}	$\nu \Sigma_{f2}$
3, CORE	1.5	0.4	0.02	0.01	0.085	0.135
4, ABS	1.5	0.4	0.02	0.01	0.13	0.135
5, REF	2.0	0.3	0.04	0	0.01	0

$$\chi_1 = 1.0, \chi_2 = 0, \nu \Sigma_{f1} = 0, \Sigma_{2 \rightarrow 1} = 0 \text{ all region}$$

A list of the complete run is shown as appendix A.

The output from this calculation is shown as appendix B.

REFERENCES

1. I. Misfeldt, Solution of the Multigroup Neutron Diffusion Equation by the Finite Element Method.
2. I. Misfeldt, Master thesis.
3. The application of the Finite Element Method to the multi-group neutron diffusion equation, Nuclear Science and Engineering 47. By L.A. Semanza et al.
4. The Finite Element Method in Engineering Science. By O.C. Zienkewicz.

APPENDIX A

A complete card deck for a calculation on the example shown page 8.

Output from this job is shown as appendix B.

```
?JOB PT/MISFI:CLASS=1:CHARGE=251402:USER=RTTHISFELUT/WORK
?PROCESSTIME=300: IRTIME=300: PRIELIMIT=5000:
?BEGIN COMPILE FROM ALGOL
?ALGOL FILE TAPL=IFRM
?DATA
$SET MERGE
?DATA INDDATA
```

FF	}	administration input	}	INPUT
1				
17.3.1976	}	control variables		
3.3.5.2.2.2.1.1.1.2.				
6.6.	}	mesh specification		
0.10.30.50				
1.1.1.	}	composition numbers		
2.2.2.				
0.10.30.50.	}	boundary conditions		
1.1.1.				
2.2.2.	}	ordinary materials		
1.2.2.2.2.				
1.5.5.5.2.	}	condensation specifications		
1.3.3.5.2.				
1.4.3.5.2.	}	control variables for the iterations		
1.1.1.1.1.				
1	}	administration input		
0.0.				
2.	}			
0.467.0.407.				
3	}			
1.5.0.4.				
0.03.0.0.02.0.005	}			
1.0.0.				
0.0.135.	}			
1.1.				
4.	}			
1.5.0.4.				
0.03.0.0.02.0.13.	}			
1.0.0.				
0.0.135.	}			
1.1.				
5.	}			
2.0.0.3.				
0.04.0.0.00.0.01.	}			
1.0.				
0.0.	}			
0.0.				
1.0.1.0.1.0.1.	}			
1.0.1.0.1.0.1.				
-1.-1.-1.-1.	}			
75.-1.-1.1.2.				
FF	}			
-1.				

?END JOB

APPENDIX B

Output from a calculation on the example shown page 8.

-1-

COMPILED: 21/04-1977. 22.25
 21/04-1977. 22.26
 PROCESSOR = 0 SEC
 ELAPSED = 5 SEC
 INPUT/OUTPUT = 0 SEC

A.E.K. PROGRAM NO. 10 PROBLEM NO. 1 17/3.1976
 FLUXHLEGNINGFR ELEMENTMETODEN

CMX	CMY	CLP	HH	NDU	Ng	NRH	KDS	NREC	M
3.	3.	5.	2.	2.	2.	1.	1.	1.	2.

MA	MY
6.	6.

XC			
0.	1.000F+01	3.000E+01	5.000E+01

FHX			
	1.	1.	1.
FACX/	2.	2.	2.

YC			
0.	1.000F+01	3.000E+01	5.000E+01

FHY			
	1.	1.	1.
FACY	2.	2.	2.

COMPOSITIONS NUMBERS

1.	2.	2.	2.	2.
1.	3.	3.	3.	3.
1.	4.	3.	3.	3.
1.	1.	1.	1.	1.

1
 EXTPL: 0. 0.

page 7 cont.

2
EXTPL: 4.67000E-01 4.67000E-01

3
D: 1.50000E+00 4.00000E-01
SS: 3.00000E-02 0. 2.00000E-02 8.50000E-02
FS: 1.00000E+00 0.
NFS: 0. 1.35000E-01
HR : 1 1.00000E+00 1.00000E+00

4
D: 1.50000E+00 4.00000E-01
SS: 3.00000E-02 0. 2.00000E-02 1.30000E-01
FS: 1.00000E+00 0.
NFS: 0. 1.35000E-01
HR : 1 1.00000E+00 1.00000E+00

5
D: 2.00000E+00 3.00000E-01
SS: 4.00000E-02 0. 4.00000E-02 1.00000E-02
FS: 1.00000E+00 0.
NFS: 0. 0.
HR : 1 0. 0.

MESIX

1.000E+00	0.	1.000E+00	0.	1.000E+00	0.	1.000E+00
-----------	----	-----------	----	-----------	----	-----------

MESIX

1.000E+00	0.	1.000E+00	0.	1.000E+00	0.	1.000E+00
-----------	----	-----------	----	-----------	----	-----------

MAXIMUM ITERATION 1
50.

EPS
1.0000E-04

EPSMV
1.0000E-01

OMEGA
1.2000E+00

-2-

21/04-1977. 22.26
 PRUCESOR = 4 SEC
 ELAPSED = 21 SEC
 INPUT/OUTPUT = 1 SEC

A.E.K. PROGRAM NO. 10 PROBLEM NO. 1 17/3.1976
 FLUXPLKLGINGER ELFHENTHTTOEN

MAXH 2916. MAXANTALITERATIONEP 50. EPS 1.0000E-04 EPSMY 1.0000E-01 OMEGA 1.2000E+00
 DANTAL 5 RTYPER 2 GANTAL 2 EANTAL 9 DANTAL 12 ANTALUBK 49 MAXHUB 625 ANTALBLK 1

IT. NR	RESIDUE	RATIO	LAMBDA	KEFF	DELTAKEFF	TOTALKILDE
1	7.0391F-01	-0.06000000	4.50387339	0.21911213	-7.80699E-01	9.8649E+01
2	9.0071F-01	-0.59341881	1.41347284	0.70747733	4.8837E-01	8.5922E+01
3	6.3546F-01	-0.60887844	1.27092998	0.78678206	7.9205E-02	9.5544E+01
4	3.4410F-01	-0.65628040	1.24784523	0.80138144	1.4599E-02	9.7368E+01
MY: -0.656280	FRSFAC: 0.603763					
5	6.5049F-02	0.26012422	1.23163916	0.81119261	1.0545E-02	9.8649E+01
6	6.0408F-02	0.60479865	1.20698252	0.82851241	1.6586E-02	1.0066E+02
7	4.4897F-02	0.76583404	1.16818730	0.84161813	1.3106E-02	1.0226E+02
8	3.8365F-02	0.77477400	1.18240903	0.84573102	1.1129E-03	1.0276E+02
9	3.3256F-02	0.84818218	1.17090095	0.85404320	8.3122E-03	1.0337E+02
10	2.8495F-02	0.81018541	1.16987375	0.85479309	7.4489E-04	1.0386E+02
11	2.3052F-02	0.65704474	1.16329243	0.85962908	4.8360E-03	1.0444E+02
12	2.0253F-02	0.80714359	1.16239254	0.86029458	6.6550E-04	1.0453E+02
13	1.5890F-02	0.85212703	1.15860164	0.86310949	8.8149E-03	1.0487E+02
14	1.3440F-02	0.81378231	1.15782383	0.86360926	5.7963E-04	1.0494E+02
15	1.0842F-02	0.64824208	1.15546132	0.86545520	1.7659E-03	1.0515E+02
16	9.5138F-03	0.62043426	1.15487357	0.86589565	4.4646E-04	1.0521E+02
17	7.4559F-03	0.84594760	1.15338089	0.86701627	1.1206E-03	1.0534E+02
18	6.4751F-03	0.82708436	1.15292799	0.86735686	3.4059E-04	1.0538E+02
19	5.1638F-03	0.84397360	1.15194595	0.86805861	7.0175E-04	1.0547E+02
20	4.1970F-03	0.83005635	1.15165025	0.86831918	2.6057E-04	1.0550E+02
21	3.4843F-03	0.84098625	1.15106760	0.86875071	4.3953E-04	1.0555E+02
MY: 0.840986	FRSFAC: 6.288883					
22	3.8045F-03	0.76560698	1.14947118	0.86996527	1.2066E-03	1.0570E+02
23	3.0052F-03	-0.90990436	1.14654962	0.87218205	2.2168E-03	1.0597E+02
24	2.2527F-03	-0.79709292	1.14682943	0.87045124	1.7308E-03	1.0576E+02
25	1.5279F-03	-0.62795846	1.14836318	0.87080465	3.5341E-04	1.0580E+02
MY: -0.627958	FRSFAC: 0.614266					
26	3.9843F-04	-0.39071867	1.14869800	0.87039929	4.0536E-04	1.0575E+02
27	2.8890F-04	0.57651044	1.14907872	0.87026240	1.3689E-04	1.0574E+02
28	1.8438F-04	0.64111460	1.14911565	0.87023443	2.7969E-05	1.0573E+02
29	1.6022F-04	0.80962135	1.14912448	0.87022774	6.6931E-06	1.0573E+02
30	1.1537F-04	0.76858433	1.14914324	0.87021354	1.4200E-05	1.0573E+02
31	9.8336F-05	0.82882590	1.14916052	0.87020045	1.3688E-05	1.0573E+02
32	7.6159F-05	0.81161009	1.14916714	0.87019544	5.0138E-06	1.0573E+02

	33	6.3541E-05	0.82420313	1.14918109	0.87018487	-1.0562E-05	1.0573E+02
MY:	0.824203	EUSFAC : 5.688383					
	34	1.1934E-04	-1.31517513	1.14920864	0.87016401	-2.0864E-05	1.0572E+02
	35	9.3564E-05	-0.64759470	1.14927801	0.87011149	-5.2218E-05	1.0572E+02
	36	5.1352E-05	-0.67054333	1.14922709	0.87015004	3.8549E-05	1.0572E+02
	37	0.7837E-05	-0.70182664	1.14924036	0.87013999	-1.0048E-05	1.0572E+02
MY:	-0.701827	EUSFAC : 0.587604					
	38	6.8107E-06	0.35387849	1.14922867	0.87014885	8.8558E-06	1.0572E+02
	39	3.8510E-06	-0.56958953	1.14922397	0.87015240	3.5542E-06	1.0572E+02
	40	3.5677E-06	0.49979640	1.14922481	0.87015177	-6.3374E-07	1.0572E+02

page 2 cont.

-3-

21/04-1977. 22.28
 PROCESSOR = 28 SEC
 ELAPSED = 126 SEC
 INPUT/OUTPUT = 8 SEC

A.E.K. PROGRAM NO. 10 PROBLEM NO. 1 17/3.1976
 FLUXBILREGNINGER ELEMENTMETODEN

ITERATIONEN ER AFSLUTTET
 DER ER UDFØRT 40 ITERATIONER
 EPSILON VAR 0.000100000000 DET MAKSIMALE ANTAL ITERATIONER 50
 DEN TOTALE FISSIONSKILDE ER 2.7853E+05

KEFF 0.8701517680
 FLUKSEN BLEV

FLUX I GRUPPE 1
 5000.: 309. 307. 285. 246. 141. 41. 23.
 4000.: 1348.1329.1278.1022. 616. 245. 41.
 3000.: 5980.5006.5650.4728.2572. 616. 141.
 2000.: 9999.9044.8625.8489.4728.1022. 246.
 1000.: 9469.9595.9002.8626.5650.1278. 285.
 500.: 8926.9134.9595.9945.5906.1329. 307.
 0.: 8688.8926.8469.9999.5980.1348. 309.
 0. 500.1000.2000.3000.4000.5000.

FLUX I GRUPPE 2
 5000.: 178. 169. 172. 122. 104. 83. -11.
 4000.: 5337.5268.5032.4117.2626.1166. 83.
 3000.: 3805.3754.3745.2950.3050.2626. 104.
 2000.: 2178.2164.2304.1672.2958.4117. 122.
 1000.: 2157.2153.2029.2304.3745.5032. 172.
 500.: 1482.1443.2153.2164.3754.5268. 169.
 0.: 1494.1482.2157.2178.3805.5337. 178.
 0. 500.1000.2000.3000.4000.5000.

```

21/04-1977.      22.28
PROCESSOR      =    28 SEC
ELAPSED        =   128 SEC
INPUT/OUTPUT   =     8 SEC

```

I GRUPPE:	1	ER DEN MAKSIMALE MIDDELFLUX:				9215			
FLUXEN FØREKOM I	OMRÅDET	X FRA	0.	TIL	10.	OG Y FRA	0.	TIL	10.
I GRUPPE:	2	ER DEN MAKSIMALE MIDDELFLUX:				4149			
FLUXEN FØREKOM I	OMRÅDET	X FRA	30.	TIL	50.	OG Y FRA	0.	TIL	10.

5000.	-+-----+-----+-----+
	: 1912 : 1458 : 335 :
3000.	-+-----+-----+-----+
	: 9181 : 7620 : 1458 :
1000.	-+-----+-----+-----+
	: 9215 : 9181 : 1912 :
0.	-+-----+-----+-----+
	0. 1000. 3000. 5000.

5000.	-	-	-	-	-	-	-	-
	:	4	1	4	:	3	2	1
3000.	-	-	-	-	-	-	-	-
	:	2	4	5	:	2	7	8
1000.	-	-	-	-	-	-	-	-
	:	1	6	8	:	2	4	5
0.	-	-	-	-	-	-	-	-

U. 1000. 3000. 5000.
DEN TILGRIPEDE FLUX FOR DELE REAKTOREN OG ALLE GRUPPER ER: 10489068

-5-

```
21/04-1977.      22.28
PROCESSOR         =    28 SEC
ELAPSED          =   130 SEC
INPUT/OUTPUT     =     8 SEC
```

A.E.K. PROGRAM NO. 10 PROBLEM NO. 1 17/3.1976
FLUXBLEIENINGEN EFFICIENTIEUDEN

REACTIUNTEGRALER:

REAKTIONSINTEGRALERNE ER NORMIERET SAA GENNEMSNITTET FOR DEN DEL AF REAKTUREN HVOR REAKTIONSINTEGRALET OVERSTIGER $\rho = 10 \cdot \max(\text{REAKTIONSINTEGRALE})$ ER 1000.

REAKTIONSGRUPPEN INTEGRAL OBER ALLE GRUPPEN. ELEMENTVIS.
REAKTIONSGRUPPEN INTEGRAL OBER ALLE GRUPPEN. ELEMENTVIS.

5000.	+	-	+	-	+	-	+	-	+
	:		:		:		:		:
3000.	+	-	+	-	+	-	+	-	+
	:	1079	:	918	:		:		:
1000.	+	-	+	-	+	-	+	-	+
	:	1611	:	1679	:		:		:
0.	+	-	+	-	+	-	+	-	+

0. 1000. 3000. 5000.

```

21/04-1977.      22.29
PROCESSOR        =   37   SEC
ELAPSED          =  216   SEC
INPUT/OUTPUT    =    9   SEC

```

OMEGA
1.5000E+00

MAXHUB AITALLUK
2401 3

-19-

MY:	0.862977	32	5.6370E-05	0.86297760	1.16109064	0.86125920	-7.7660E-06	3.0018E+05
				FNSIAC :	7.298023			
		33	4.9971E-05	0.53009140	1.16116711	0.86120248	-5.6717E-05	3.0016E+05
		34	3.7089E-05	-0.73125440	1.16118101	0.86119217	-1.0310E-05	3.0016E+05
		35	3.2021E-05	-0.80644860	1.16116655	0.86120290	1.0729E-05	3.0016E+05
MY:	-0.629627	36	1.4191E-05	-0.62962657	1.16116301	0.86120552	2.6207E-05	3.0017E+05
				FNSIAC :	0.613638			
		37	1.0158E-05	0.50530331	1.16116014	0.86120766	2.1327E-06	3.0017E+05
		38	7.2458E-06	0.72444582	1.16115737	0.86120970	2.0490E-06	3.0017E+05
		39	3.6702E-06	0.74054001	1.16115552	0.86121108	1.3764E-06	3.0017E+05

page 6 cont.

ITERATIONEN ER AFSLUTTET
DER ER UDLEGT 39 ITERATIONER
EPSILON VAR 0.0001000000000 DET MAXIMALE ANTAL ITERATIONER 75
DEN TOTALE FISSIONSKILDE ER 2.5505E+05

-21-

[illegible]

-8-

21/04-1977. 22.32
 PROCESSION = 107 SEC
 ELAPSEU = 387 SEC
 INPUT/OUTPUT = 26 SEC

A.L.P. PROGRAM NO. 10 REWIND NO. 1 17/3.1976
 FLUXFELIGHEID LEPENTAFLEED

I GROEP 1 1 EF DEN MAXIMALE MIDDELFLOX: 9846
 FLUXF: TOEGANG I HAALET X FRA 10. TIL 20. OG Y FRA 0. TIL 5.
 I GROEP 2 2 EF DEN MAXIMALE MIDDELFLOX: 5323
 FLUXF: TOEGANG I HAALET X FRA 30. TIL 40. OG Y FRA 0. TIL 5.

MIDDELFLOX I GROEP 1

5000.	1	614	1	595	1	528	1	370	1	187	1	63
4000.	1	2759	1	2662	1	2399	1	1687	1	696	1	187
3000.	1	7911	1	7137	1	7031	1	5050	1	1687	1	370
2000.	1	5066	1	0822	1	9366	1	7031	1	2399	1	528
1000.	1	5261	1	9556	1	9622	1	7737	1	2602	1	595
500.	1	6666	1	9282	1	9046	1	7911	1	2759	1	614
0.												

0. 500. 1000. 2000. 3000. 4000. 5000.

MIDDELFLOX I GROEP 2

5000.	1	2281	1	2211	1	1968	1	1836	1	809	1	271
4000.	1	5323	1	5169	1	4616	1	3444	1	2295	1	809
3000.	1	2352	1	2298	1	2099	1	1140	1	3444	1	1836
2000.	1	2219	1	2228	1	2195	1	2099	1	4616	1	1968
1000.	1	1567	1	1765	1	2270	1	2299	1	5169	1	2211
500.	1	1399	1	1568	1	2219	1	2352	1	5323	1	2281
0.												

0. 500 1000. 2000. 3000. 4000. 5000.
DEN INTEGRENERE FLUX FOR HELE REAKTOREN OG ALLE GRUPPER ER:

15177199

page 8 cont.

```

-9-      21/04-1977.      22.32
          PRUCESSOR      =      108 SEC
          ELAPSED      =      388 SEC
          INPUT/OUTPUT =      26 SEC

```

A.E.K. PROGRAM NO. 10 PROBLEM NO. 1 17/3.1976
FLUXING/CLIPPING ELEPHANTRETSODEN

REACTANTS INTEGRAL:

REAKTIONSINTEGRALERNE ER NORMALISERT SÅA GLIENSNITTET FÜR DEN DEL AF REAKTUREN HVOR REAKTIONSINTEGRALET OVERSTIGER $9 \cdot 10 \cdot \max(\text{REAKTIONSINTEGRALE})$ ER 1000.

REAKTIONSGRUPPEN, ELEMENTAR-
REAKTIONSGRUPPEN NR. 1

5000.	-+-----+	-+-----+	-+-----+
	0	0	(
3000.	-+-----+	-+-----+	-+-----+
	1067	198	0
1000.	-+-----+	-+-----+	-+-----+
	1059	1067	0
0.	-+-----+	-+-----+	-+-----+
	0.	1000.	3000.
			5000.

```

TIME USED FOR THIS PROBLEM
PROCESSOR      =    108 SEC
ELAPSED       =    309 SEC
INPUT/OUTPUT   =     26 SEC

```

APPENDIX C

Example II

A complete card deck for a calculation with boundary condition (2), gamma matrix.

Output corresponding to this run is shown in appendix D.

```
?JOB RT/MISFT;CLASS=1;CHARGE=251002;USER=RTMISFELDT/WORK
?PROCESSTIME=300; IUTIME=300; PRINTLIMIT=5000;
?BEGIN COI PILE FENB ALGOL
?ALGOL FILE TAPE=1ENB
?DATA
$SET HLHGL
?DATA INDATA
FF
2
22,3,1977,
2.2,4.2,1,2,1,0,0,1,
8.8,
0.7,8,
7.1,
2.2,
0.7,8,
7.1,
2.2,
1.2,2,2,
1.3,3,2,
1.4,3,2,
1.1,1,1,
1,
0.0,
2,
0.2,0,-0.1,0.05,
3,
2.0,3,
0.03,0,0.03,0.01,
1.0,
0.0,
0.0,
4,
1.0,4,
0.02,0,0.01,0.01,
1.0,
0.0,05,
0.0,02,
75,-1,-1,1.5,
FF
-1,
?END JOB
```

APPENDIX D

Output for example II corresponding to the card deck shown
in appendix C.

-1-

A.E.K. PROGRAM NO. 10 PROBLEM NO. 2 22/3.1977
FLUXPHILGIRGIR ELIDENTITODEN

CHY CUY FCP LH NDB NO DFF KDS I-REC M
2. 2. 4. 2. 1. 2. 1. 0. 0. 1.

1. BY
1. 6.

YC

0. 7.000E+00 0.000E+00

FEX

7. 1.

FAC. 7

2. 2.

YC

0. 7.000E+00 0.000E+00

FEX

7. 1.

FAC. 7

2. 2.

COMPOSITIONS TO BE EPS

1. 2. 2. 2.
1. 3. 3. 2.
1. 4. 3. 2.
1. 1. 1. 1.

1
FATHER C. C.

2
 GANNA: 2.00000E-01 0. -1.00000E-01 5.00000E-02

3
 DS: 2.00000E+00 3.00000E-01
 SS: 3.00000E-02 0. 3.00000E-02 1.00000E-02
 FS: 1.00000E+00 0.
 NFS: 0. 0.
 NR 11 0. 0.

4
 DS: 1.00000E+00 4.00000E-01
 SS: 2.00000E-02 0. 1.00000E-02 1.00000E-02
 FS: 1.00000E+00 0.
 NFS: 0. 5.00000E-02
 NR 11 0. 2.00000E-02

MAXIMUM INITIAL TIME: 75. LPS 1.00000E+00 EFSHY 1.00000E-01 DMLCA 1.50000E+00

page 1 cont.

21/04-1977. 22.32
PROCESSOR ■
TRANSFER ■
INPUT/OUTPUT ■

TAXI 1024. JAVANTALITERATION 75.				LPS 1.0000E-06		EPSKY 1.0000E-01		OMEGA 1.5000E+00	
PANTAL	PTYPE	CANTAL	ESNTAL	PANTAL	ANTALBLK	MAXHDD	ANTALBLK		
4	2	1	64	32	81	625	1		

10. 11	12. 11	13. 11	14. 11	15. 11	16. 11	17. 11	18. 11	19. 11	20. 11	21. 11	22. 11	23. 11	24. 11	25. 11	26. 11	27. 11	28. 11	29. 11	30. 11	31. 11	32. 11	33. 11	34. 11	35. 11	36. 11	37. 11	38. 11	39. 11	40. 11	41. 11	42. 11	43. 11	44. 11	45. 11	46. 11	47. 11	48. 11	49. 11	50. 11	51. 11	52. 11	53. 11	54. 11	55. 11	56. 11	57. 11	58. 11	59. 11	60. 11	61. 11	62. 11	63. 11	64. 11	65. 11	66. 11	67. 11	68. 11	69. 11	70. 11	71. 11	72. 11	73. 11	74. 11	75. 11	76. 11	77. 11	78. 11	79. 11	80. 11	81. 11	82. 11	83. 11	84. 11	85. 11	86. 11	87. 11	88. 11	89. 11	90. 11	91. 11	92. 11	93. 11	94. 11	95. 11	96. 11	97. 11	98. 11	99. 11	100. 11	101. 11	102. 11	103. 11	104. 11	105. 11	106. 11	107. 11	108. 11	109. 11	110. 11	111. 11	112. 11	113. 11	114. 11	115. 11	116. 11	117. 11	118. 11	119. 11	120. 11	121. 11	122. 11	123. 11	124. 11	125. 11	126. 11	127. 11	128. 11	129. 11	130. 11	131. 11	132. 11	133. 11	134. 11	135. 11	136. 11	137. 11	138. 11	139. 11	140. 11	141. 11	142. 11	143. 11	144. 11	145. 11	146. 11	147. 11	148. 11	149. 11	150. 11	151. 11	152. 11	153. 11	154. 11	155. 11	156. 11	157. 11	158. 11	159. 11	160. 11	161. 11	162. 11	163. 11	164. 11	165. 11	166. 11	167. 11	168. 11	169. 11	170. 11	171. 11	172. 11	173. 11	174. 11	175. 11	176. 11	177. 11	178. 11	179. 11	180. 11	181. 11	182. 11	183. 11	184. 11	185. 11	186. 11	187. 11	188. 11	189. 11	190. 11	191. 11	192. 11	193. 11	194. 11	195. 11	196. 11	197. 11	198. 11	199. 11	200. 11	201. 11	202. 11	203. 11	204. 11	205. 11	206. 11	207. 11	208. 11	209. 11	210. 11	211. 11	212. 11	213. 11	214. 11	215. 11	216. 11	217. 11	218. 11	219. 11	220. 11	221. 11	222. 11	223. 11	224. 11	225. 11	226. 11	227. 11	228. 11	229. 11	230. 11	231. 11	232. 11	233. 11	234. 11	235. 11	236. 11	237. 11	238. 11	239. 11	240. 11	241. 11	242. 11	243. 11	244. 11	245. 11	246. 11	247. 11	248. 11	249. 11	250. 11	251. 11	252. 11	253. 11	254. 11	255. 11	256. 11	257. 11	258. 11	259. 11	260. 11	261. 11	262. 11	263. 11	264. 11	265. 11	266. 11	267. 11	268. 11	269. 11	270. 11	271. 11	272. 11	273. 11	274. 11	275. 11	276. 11	277. 11	278. 11	279. 11	280. 11	281. 11	282. 11	283. 11	284. 11	285. 11	286. 11	287. 11	288. 11	289. 11	290. 11	291. 11	292. 11	293. 11	294. 11	295. 11	296. 11	297. 11	298. 11	299. 11	300. 11	301. 11	302. 11	303. 11	304. 11	305. 11	306. 11	307. 11	308. 11	309. 11	310. 11	311. 11	312. 11	313. 11	314. 11	315. 11	316. 11	317. 11	318. 11	319. 11	320. 11	321. 11	322. 11	323. 11	324. 11	325. 11	326. 11	327. 11	328. 11	329. 11	330. 11	331. 11
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33	2.3692F-04	0.99023407	1.00271865	0.92360098	1.7421E-04	2.2628E+00
34	2.2646F-04	0.9792045A	1.00249824	0.923378403	1.06006E-04	2.2633E+00
35	2.2614F-04	0.97924162	1.00226814	0.923398544	1.9640E-04	2.2638E+00
36	2.2619F-04	0.96066417	1.00203329	0.9224185598	2.0034E-04	2.2644E+00
37	2.261490F-04	0.95429676	1.00179781	0.9224330715	2.0117E-04	2.2647E+00
38	2.261003F-04	0.94443462	1.00156472	0.922458637	1.9422E-04	2.2652E+00
39	2.260648F-04	0.94535339	1.00133640	0.922478159	1.9522E-04	2.2657E+00
40	2.260104F-04	0.94180678	1.00111468	0.922497125	1.8966E-04	2.2662E+00
NY: 0.941807	FKS: AC: 17.144131					
41	4.5647F-04	0.75150320	1.07745331	0.922811446	3.1432E-03	2.2739E+00
42	3.0904F-04	0.65696127	1.07738122	0.922817656	6.2104E-03	2.2740E+00
43	2.0953F-04	0.86782146	1.07732074	0.922822867	5.2109E-03	2.2742E+00
44	1.3720F-04	0.93514711	1.07727245	0.922827028	4.1605E-03	2.2743E+00
45	1.0274F-04	0.80189753	1.07723954	0.922829664	2.8335E-03	2.2743E+00
46	0.8779F-05	0.85512709	1.07722094	0.922831463	1.5995E-03	2.2744E+00
47	7.7477F-05	0.87606048	1.07721488	0.922831989	3.2222E-03	2.2744E+00
48	6.4420F-05	0.92606723	1.07721599	0.922831635	3.5433E-03	2.2744E+00
49	5.5108F-05	0.92143062	1.07723133	0.922830567	1.0078E-03	2.2743E+00
50	4.6670F-05	0.96074077	1.07724923	0.922829029	1.5382E-03	2.2743E+00
51	4.1744F-05	1.02184114	1.07727230	0.922827041	1.9660E-03	2.2743E+00
52	3.6407F-05	1.00029816	1.07729986	0.922824666	3.7499E-03	2.2742E+00
53	3.5951F-05	0.96368765	1.07733062	0.922822016	6.5033E-03	2.2741E+00
54	3.5093F-05	0.97283342	1.07736348	0.922819185	8.8311E-03	2.2741E+00
55	3.3913F-05	0.96464538	1.07739759	0.922816246	2.9387E-03	2.2740E+00
56	3.2721F-05	0.95841353	1.07743225	0.922813261	4.9655E-03	2.2739E+00
57	3.1804F-05	0.95254364	1.07746690	0.922810276	6.9647E-03	2.2739E+00
58	3.1002F-05	0.94785676	1.07750110	0.922807330	8.9458E-03	2.2738E+00
59	3.0447F-05	0.94395331	1.07753452	0.922804452	1.0878E-03	2.2737E+00
60	2.9538F-05	0.94050083	1.07756690	0.922801663	2.7691E-03	2.2736E+00
NY: 0.940580	FKS: AC: 16.829472					
61	6.7509F-05	0.74114431	1.07309162	0.92756395	4.5168E-04	2.2725E+00
62	4.4020F-05	0.61583084	1.07310165	0.92755632	6.6258E-04	2.2725E+00
63	2.7669F-05	0.86703666	1.07311012	0.92754903	7.2425E-04	2.2725E+00
64	1.9312F-05	0.88623485	1.07311695	0.92754316	5.8735E-04	2.2725E+00
65	1.4342F-05	0.96252362	1.07312165	0.92753911	4.0455E-04	2.2725E+00
66	1.2369F-05	0.93533032	1.07313436	0.92753678	3.3279E-04	2.2725E+00
67	1.1095F-05	0.97616519	1.07313533	0.92753595	3.3619E-04	2.2725E+00
68	8.9615F-06	0.92256270	1.07312487	0.92753534	3.9293E-04	2.2725E+00
69	7.6636F-06	0.91884529	1.07312326	0.92753773	1.3683E-04	2.2725E+00
70	6.5213F-06	0.90790624	1.07312088	0.92753977	2.0443E-04	2.2725E+00
71	5.7928F-06	1.02404059	1.07311777	0.92754245	2.6757E-04	2.2725E+00

page 2 cont.

```

ITERATIONEN ER AFSLUTTET
DER ER GÅET      71 ITERATIONER
EPSILO VAR 0.000100000000 DET MÅKSIMALE ANTAL ITERATIONER    75
DET TILFÆLLE PÅSSIGESIDE ER      2.2196E+04

```

FLUX	GROUP	5234	5197	5085	4899	4640	4314	3942	3572	3224
8000	1	5837	5790	5672	5469	5177	4809	4373	3917	3572
7000	1	7061	7012	6864	6614	6254	5789	5183	4373	3942
6000	1	6025	7079	7000	7011	7094	6531	5789	4809	4314
5000	1	5773	5703	4519	4116	7724	7094	6259	5177	4641
4000	1	9322	9252	9051	8703	8196	7511	6515	5466	4894
3000	1	4704	4632	4417	4051	3519	7300	6304	5672	5080
2000	1	9926	9853	9632	9255	8703	7470	7012	5796	5197
1000	1	4099	4026	4793	4322	3770	3027	7062	5837	5235
0	1	0	1000	2000	3000	4000	5000	6000	7000	8000

FLUX 1	GROUP 2
8000.:	9329.9336.9495.9345.9148.8913.8572.8417.8145.
7000.:	9375.9371.9301.9192.9052.8896.8741.8600.8417.
6000.:	9214.9196.9135.9052.8953.8861.8736.8641.8572.
5000.:	9127.9113.9073.9015.8944.8881.8801.8726.8618.
4000.:	9097.9086.9057.9015.8973.8942.8913.8852.8749.
3000.:	9097.9086.9057.9033.9015.9015.9002.9142.9346.
2000.:	9111.9104.9037.9033.9057.9071.9143.9301.9495.
1000.:	9123.9110.9104.9062.9066.9113.9199.9371.9598.
0.:	9123.9123.9111.9097.9097.9127.9215.9345.9620.
	0.1000.2000.3000.4000.5000.6000.7000.8000.

-4-

21/04-1977. 22.36
 PRCESSOR = 96 SEC
 ELAPSEU = 195 SEC
 INPUT/OUTPUT = 13 SEC

A.E.K. PROGRAM NO. 10 PROBLEM NO. 2 22/3.1977
 FLUXBERECHNUNG LIFEBEHTNETZDEK

I GRUPE: 1 ER DEN MAXIMALE MIDDLEFLUX: 9926
 FLUXEN FOLGEN I BEZAHLT X FRA 0. TIL 1. OG Y FRA 0. TIL 1.
 I GRUPE: 2 ER DEN MAXIMALE MIDDLEFLUX: 9494
 FLUXEN FOLGEN I BEZAHLT X FRA 7. TIL 8. OG Y FRA 0. TIL 1.

MIDDLEFLUX I GRUPE 1

8000.	5516	5638	5241	5046	4735	4360	3951	3572
7000.	6627	6336	6154	5879	5508	5040	4463	3751
6000.	7518	7412	7197	6869	6418	5824	5040	4360
5000.	8369	8249	8001	7632	7112	6418	5500	4735
4000.	9014	8863	8617	806	7632	6869	5879	5046
3000.	9478	9339	9055	8617	8007	7197	6154	5281
2000.	9778	9633	9339	8883	8249	7412	6336	5438
1000.	9926	9778	9478	9014	8369	7518	6427	5516
0.								

0. 1000. 2000. 3000. 4000. 5000. 6000. 7000. 8000.

MIDDLEFLUX I GRUPE 2

8000.	9494	9439	9333	9184	9003	8807	8608	8395
7000.	9298	9253	9175	9067	8943	8818	8715	8606
6000.	9163	9131	9073	8997	8915	8847	8818	8807
5000.	9106	9062	9046	8988	8939	8915	8943	9003
4000.	9092	9075	9085	9011	8988	8997	9067	9184

0. 1000. 2000. 3000. 4000. 5000. 6000. 7000. 8000. 1022746
DEN TILGEGNEDE FLUX FOR HJELP REAKTORER OG ALLE GRUPPER ER:

-32-

-5-

21/04-1977. 22.36
 PRØCESOR = 97 SEC
 ELAPSED = 196 SEC
 INPUT/OUTPUT = 13 SEC

A.E.K. PROCPAD NO. 10 PROBLEM NO. 2 22/3.1977
 FLUXKORREKTIONER I ELEMENTETIDEN

REAKTIONSGRUPPETER:

REAKTIONSGRUPPETERNE ER INDDELLET I 10 GRUPPER. SAM GEMIDTET FOR DEN DEL AF REAKTØREN
 HVOR REAKTIONSGRUPPETERNE OVERSTIGER $\rho = 10 \cdot \max(\text{REAKTIONSGRUPPETER})$ ER 1000.

REAKTIONSGRUPPETER OVER ALLE GRUPPER. ELEMENTVIS.
 REAKTIONSGRUPPETER NO. 1

8000.	1	6	1	9	1	0	1	0	1	0	1	0	1	0	1	0	1
7000.	1	1026	1	1001	1	1013	1	1001	1	937	1	973	1	962	1	0	1
6000.	1	1011	1	1008	1	1002	1	993	1	934	1	977	1	973	1	0	1
5000.	1	1005	1	1002	1	997	1	992	1	987	1	984	1	987	1	0	1
4000.	1	1000	1	1002	1	998	1	995	1	992	1	993	1	1001	1	0	1
3000.	1	1005	1	1003	1	1001	1	994	1	996	1	1002	1	1013	1	0	1
2000.	1	1006	1	1005	1	1003	1	1002	1	1002	1	1004	1	1021	1	0	1
1000.	1	1007	1	1006	1	1005	1	1004	1	1005	1	1011	1	1026	1	0	1
0.	1	1007	1	1006	1	1005	1	1004	1	1005	1	1011	1	1026	1	0	1

0. 1000. 2000. 3000. 4000. 5000. 6000. 7000. 8000.

TIME USED FOR THIS PROBLEM
 PRØCESOR = 97 SEC
 ELAPSED = 196 SEC
 INPUT/OUTPUT = 13 SEC